Bipolar Electrochemistry with Organic Single Crystals for Wireless Synthesis of Metal–Organic Janus Objects and Asymmetric Photovoltage Generation

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ABSTRACT: Bipolar electrochemistry has recently emerged as a very unique method to address conducting particles in a wireless manner. The technique is often applied to the fabrication of Janus particles; however the chemical nature of the bipolar electrode has been essentially limited to carbon- or metal-based materials. Here, we report for the first time the use of conducting organic single crystals as bipolar electrodes for the preparation of a new generation of Janus objects. Fabre and Bechgaard salts involving respectively tetrathia- and tetraselenafulvalene were selected for proof-of-concept experiments. Such an approach allows to preserve the integrity of these fragile substrates because it necessitates neither electronic wiring nor mechanical contact. The site-selective electrodeposition of copper is successfully achieved, leading thus to a new metal–organic Janus structure. Subsequently, asymmetric generation of photovoltage under illumination is achieved due to the anisotropic presence of copper, making this approach interesting for the design of novel hybrid objects with applications in organic electronics or photocatalysis.

INTRODUCTION

Janus particles are the focus of major ongoing research studies because this class of asymmetric materials exhibits unprecedented anisotropic properties with many possible applications, especially for electronics, advanced spectroscopy, and optics. Several synthetic approaches have been proposed for the preparation of Janus objects whose common feature being indeed the necessity to break the symmetry of a chemical system.1–7

Among the various fabrication methods reported so far, bipolar electrochemistry (BPE) has recently emerged as a very unique approach.8–11 BPE is already known for several decades and has been particularly used in the context of electrolysis and corrosion science.12–16 In recent years, however, BPE has been reinvestigated and is nowadays a very hot topic with a much broader range of applications, especially in materials science and analytical chemistry.17–24 BPE is an unconventional approach compared to a classic electrochemical setup where the electrochemical reactions take place at the surface of two different electrodes, namely the working and auxiliary electrode both connected to a potentiostat. In the case of BPE, an electric field is applied across an electrolyte solution where a conducting object is immersed and behaves as a floating electrode. The driving force is therefore the polarization potential established between the solution and the object, allowing thus to promote oxidation and reduction reactions at the opposite poles of the same object (Figure 1). BPE can be considered as an electrochemical break of symmetry of the corresponding bipolar electrode which behaves simultaneously as an anode and a cathode. Also, the bipolar electrode is...
addressed electrochemically in a wireless fashion because there is no direct connection with the power supply. Finally, one of the key advantages of BPE is that it does not directly involve a 2-dimensional interface to break the symmetry, providing a powerful tool for the bulk (i.e., 3D) fabrication or activation of Janus objects.\textsuperscript{25,26} It is also noteworthy that BPE can be used to generate surface anisotropy based on the gradient of polarization potential alongside the bipolar electrode. This has been exemplified by several contributions showing the variation of chemical composition, morphology, or thickness and therefore of the resulting physical, optical, or spectroscopic properties.\textsuperscript{7,27–37}

So far, BPE has proven to be very versatile for site-selective wireless electrodeposition by tuning the nature of the chemical species involved in the bipolar reactions at the extremities of a conducting object. For instance, the reduction of a metal precursor to form the corresponding metal,\textsuperscript{38–40} the oxidation of a monomer to form a polymer layer,\textsuperscript{41} the electrografting of an organic layer,\textsuperscript{42} pH-induced local precipitation,\textsuperscript{43} and the deposition of various inorganic structures were reported.\textsuperscript{44,45} However, all these experiments were performed on bipolar electrodes made out of either metal or carbon allotropes, thus exhibiting a high electrical conductivity.\textsuperscript{46–48} The reason is obviously because BPE is favored when the bipolar electrode is more conductive than the surrounding media due to the competition between the Faradic current flowing through the bipolar electrode and the ionic current occurring across the electrolyte. As a matter of fact, varying the chemical nature of the bipolar electrode has not been widely investigated even if BPE experiments with other inorganic bipolar electrodes such as tin-doped indium oxide (ITO) surfaces or TiO\textsubscript{2} tubes were reported.\textsuperscript{49} It is also noteworthy that organic thin layers based on conductive polymers were reported to behave as bipolar surfaces.\textsuperscript{28,30,32}

In order to extend the application range of Janus particles prepared by BPE, we report in the present contribution for the first time the use of organic single crystals as bipolar electrodes. We selected Fabre and Bechgaard salts, which are well-known organic conductor structures based on tetramethyltetrafulvalene (TMTTF) and tetramethyltetraselenafulvalene (TMTSF) as donors (Scheme 1) and hexafluorophosphate as counterion.\textsuperscript{50} These materials have been identified as possible bipolar electrode candidates because they exhibit a suitable conductivity and have also interesting potential applications in organic electronics or photocatalysis. It is also noteworthy that such organic crystals need to be handled carefully because they are relatively fragile and can therefore be irreversibly damaged during conventional electrodeposition experiments. Here, the key advantage of BPE is indeed the possibility to modify such crystals without any electronic or mechanical contact.

\begin{figure}
\centering
\includegraphics[width=0.5\textwidth]{Scheme1.png}
\caption{Chemical Structure of Tetramethyltetrafulvalene (TMTTF) and Tetramethyltetraselenafulvalene (TMTSF)}
\end{figure}

\section*{RESULTS AND DISCUSSION}

\subsection*{Chemicals.}
(TMTTF)\textsubscript{2}PF\textsubscript{6} and (TMTSF)\textsubscript{2}PF\textsubscript{6} crystals used in this study were classically prepared by electrocrystallization as described in the literature.\textsuperscript{51,52} All aqueous solutions were prepared with Milli-Q grade water (Millipore, resistivity 18.2 MΩ cm at 25 °C). Copper sulfate (CuSO\textsubscript{4}, anhydrous, 98% purity) was purchased from Alfa Aesar. Hydroquinone (HOC\textsubscript{6}H\textsubscript{4}OH, abbreviated H\textsubscript{2}Q, 99.5% minimal purity) was purchased from Sigma-Aldrich. Dichloromethane (CH\textsubscript{2}Cl\textsubscript{2}, chromasolv, ≥99.8%) and tetrabutylammonium hexafluorophosphate (Bu\textsubscript{4}NPF\textsubscript{6}) for electrochemical analysis, ≥99.0% were purchased from Sigma-Aldrich (supplier, grade). Tetramethyltetraselenafulvalene (TMTSF, 97%) was purchased from Acros Organics.

\subsection*{Bipolar Electrochemistry.}
Bipolar electrodeposition experiments were performed in a homemade cell comprising two plastic cuvettes connected by a glass capillary. The studied organic crystals were positioned inside the capillary prior to BPE experiments. The feeder electrodes were gold plates (10 × 2.5 mm) purchased from ACM (78600 Villiers, France). The distance between both feeder electrodes is ∼50 mm, and they are connected to an external power supply (Keithley 6157 electrometer).

\subsection*{Microscopy.}
Organic crystals were characterized by using an optical stereomicroscope (Leica MSV266) and a scanning electron microscope (Hitachi, TM-1000).

\subsection*{Cyclic Voltammetry.}
Cyclic voltammetry (CV) experiments were carried out in a conventional electrochemical cell comprising a gold working electrode (ϕ = 2.0 mm), a Pt mesh counter electrode, and a silver wire pseudoreference connected to a potentiostat (Autolab, PGSTAT12). The working electrode was carefully polished with alumina and rinsed before each experiment, and the CVs were recorded in CH\textsubscript{2}Cl\textsubscript{2} containing 1 mM TMTTF and 0.1 M Bu\textsubscript{4}NPF\textsubscript{6} at room temperature. CVs were measured at different scan rates ranging from 0.01 to 1.00 V s\textsuperscript{-1}.

\subsection*{Photovoltage Measurements.}
Photovoltage measurements were performed with two carbon Scotch tape electrodes (SPI Supplies, 8 mm width) with ∼1 mm spacing and connected to a high-precision voltmeter (Keithley 2001 multimeter). A LabVIEW script (version 9.0) was written and used to record the photovoltage signal. Organic crystals were carefully positioned between both electrodes prior to measurements. A commercial red laser beam (λ = 635 nm) was used for the illumination of the extremities of these crystals.
The electrochemical reactions that need to be coupled across the bipolar electrode are therefore cupric ion reduction at the cathode side (eq 1) and water oxidation at the anodic pole (eq 2):

$$\text{Cu}^{2+} + 2e^- \rightarrow \text{Cu}^0 \quad (E^\circ_1 = 0.34 \text{ V vs NHE}) \quad (1)$$

$$2\text{H}_2\text{O} \rightarrow \text{O}_2 + 4\text{H}^+ + 4e^- \quad (E^\circ_2 = 1.23 \text{ V vs NHE}) \quad (2)$$

It means that in order to achieve BPE-induced copper deposition, the minimal polarization potential difference between the two poles of the bipolar electrode should be $1.23 \text{ V} - 0.34 \text{ V} = 0.89 \text{ V}$. BPE experiments were conducted in an electrochemical cell built with two reservoirs, linked together by a glass capillary, each side containing a gold feeder electrode connected to an external power supply. The organic crystal is positioned in the capillary, and the whole cell is filled with a 10 mM aqueous solution of CuSO$_4$. The applied electric field should be at least 8.9 V cm$^{-1}$ if considering a 1 mm long crystal. Also, the distance between both feeder electrodes was fixed at 5 cm, meaning that under these experimental conditions a potential of at least $\sim 45 \text{ V}$ should be imposed by the external power supply in order to achieve the necessary electric field strength.

Bipolar electrodeposition was conducted on both (TMTTF)$_2$PF$_6$ and (TMTSF)$_2$PF$_6$ crystals, and the corresponding experiments were monitored in situ by optical microscopy while applying the electric field (Figures 2a and 2b).

From these representative optical images, one can easily identify the organic crystal positioned inside the capillary and also a marked coloration of the solution in yellow for TMTTF (Figure 2a) and in red for TMTSF (Figure 2b) with a maximum intensity located at the left side of the bipolar electrode, being actually the anodic pole. These colored compounds seem to migrate towards the feeder cathode, revealing that they are indeed positively charged molecules. Both sides of the crystals were also characterized by scanning electron microscopy (SEM) as shown in Figures 2c–f. These images show that the electrodeposition of Cu as discrete clusters (size <10 μm) does occur at the cathodic extremity of both crystals (Figures 2d and 2f). On the other hand, the anodic pole of the crystal was clearly damaged during BPE, and the corresponding images show an extended portion of the crystals where electro-induced dissolution seems to take place (Figures 2c and 2e). Therefore, an oxidative degradation of both TMTTF- and TMTSF-based crystals was observed under BPE conditions, encouraging us to further investigate the electrochemical characterization of these two donors. It is noteworthy that absolutely no degradation was observed without applying the electric field, meaning that both crystals are electrochemically stable at open-circuit potential.

The electrochemical behavior of the donor moieties was studied by cyclic voltammetry (CV) recorded in an aprotic organic solvent because of their low solubility in water. Both CV experiments were comparable, and a typical CV recorded at several scan rates is presented in Figure 3. These data were obtained from a 1 mM TMTSF solution in CH$_2$Cl$_2$ containing 0.1 M Bu$_4$NPF$_6$ as supporting electrolyte.

The cyclic voltammogram features two well-defined reversible processes corresponding to two stepwise monoelectronic oxidations of the tetraselenafulvalene moiety. The first oxidation is centered at 0.36 V versus a silver pseudoreference and corresponds to the removal of an electron from TMTSF to form a radical cation, whereas the second one occurs at 0.83 V and is assigned to the formation of a dication:

$$\text{TMTSF} \rightarrow \text{TMTSF}^{++} + e^- \quad (E^\circ_2 \sim 0.56 \text{ V vs NHE}) \quad (3)$$
These organic crystals (Figure 2). Indeed, the reduction of Cu^{2+} This is why our TMTTF and TMTSF are much easier to oxidize than water.

In aqueous phase with the MB u4NPF6 as supporting electrolyte recorded with a gold working electrode at various scan rates (\( \nu = 0.05, 0.10, 0.250, 0.50, 0.75, \) and 1.00 V s\(^{-1}\), respectively). The counter electrode was a platinum mesh, and a silver wire was used as pseudoreference.

\[
\text{TMTSF}^{+} \rightarrow \text{TMTSF}^{2+} + e^- \quad (E^0 \sim 1.03 \text{ V vs NHE})
\] (4)

These values are consistent with previously reported data (0.33 and 0.70 V versus SCE for TMTTF in CH\(_2\)CN).\(^{56}\) Even if the BPE experiments were performed in aqueous phase with the Bechgaard salts, whereas the electrochemical characterization of the donors was carried out in an organic solvent, it is clear that TMTTF and TMTSF are much easier to oxidize than water. This is why our first set of BPE tests led to a degradation of these organic crystals (Figure 2). Indeed, the reduction of Cu^{2+} at the cathodic pole is coupled with the oxidation of the crystal at the anodic side. It is also noteworthy that the changes in color observed with optical microscopy (Figure 2a,b) are attributed to the electro-oxidation of TMTTF\(^{0/+}\) or TMTSF\(^{0/+}\) into the corresponding dications as already observed for example with TTF by in situ visible absorption spectroelectrochemistry.\(^{57}\)

On the basis of these new data, another set of BPE experiments was carried out by adding to the electrolyte a sacrificial coreactant which is easier to oxidize than the donor in the organic crystal. Here, it is important to remind that the donor molecules in the crystal structures are overlapping symmetrically giving rise to a formally 1/2 filled band in the dimerized situation. For that purpose, hydroquinone (H\(_2\)Q) was selected because it exhibits appropriate redox properties with a formal oxidation potential less positive than the oxidation of the fulvalene donor into the dicationic forms TMTTF\(^{2+}\) or TMTSF\(^{2+}\), respectively:

\[
H_2Q \rightarrow Q + 2H^{2+} + 2e^- \quad (E^0_5 \sim 0.70 \text{ V vs NHE})
\] (5)

As a consequence, the two redox half-reactions involved in the BPE electrodeposition will become the ones reported in eqs 1 and 5, and therefore the necessary electric field will be much lower than previously.

Therefore, a new series of BPE tests was conducted by applying an electric field typically lower than 3–4 V cm\(^{-1}\), and the corresponding modified crystals (length \( \geq 1 \text{ mm} \) ) were characterized by SEM imaging (Figure 4).

As one can see in the corresponding micrographs, such an approach which takes advantage of a sacrificial redox coreactant was successful. The cathodic edge was indeed correctly modified with copper (Figures 4b and 4d) while the anodic side remained undamaged (Figures 4a and 4c). In the latter cases, an electric field of 2.6 V cm\(^{-1}\) was applied and resulted in a metal deposit that is well-focused at the very edge of the organic crystals. A large number of asymmetric objects of various lengths were prepared according to the same procedure, and as expected, a longer deposition zone was observed when using the longest crystals (2.5–3 mm) such as the one shown in Figure 5a. These long single crystals were then selected for further electrical characterization as they are easier to handle.

Photovoltage measurements were recorded to demonstrate the influence of the localized deposit on the physical properties of the newly prepared metal/organic hybrid material. To do so, a very simple experimental setup was chosen where the organic crystal is positioned between two carbon-tape stripes connected to a voltmeter (Figure 5b). The open-circuit voltage was then measured while illuminating either one side or the other of the asymmetric materials with a red laser (\( \lambda = 650 \text{ nm} \). A representative time evolution of the photovoltage generated across a modified (TMTSF)\(_2\)PF\(_6\) crystal as a function of the light excitation is shown in Figure 5c. Here, the geometry of the device is similar to a metal/semiconductor/metal (MSM) photodetector architecture. It means that in this configuration two metal/semiconductor Schottky contacts are positioned in front of each other, and therefore the photovoltage polarity depends on the illumination side.\(^{58}\) Indeed, the intensity of the photovoltage was found to be asymmetric with a different amplitude depending on whether the excitation was focused on the modified versus nonmodified side of the crystal. This is assigned to the different nature of the contact (and corresponding work functions) between the right and left sides with or without the presence of copper. A control experiment was also recorded with a pristine crystal. In this

Figure 4. Scanning electron micrographs of the left (a) and right (b) extremities of a (TMTTF)\(_2\)PF\(_6\) organic crystal where the anode side remains intact (due to the addition of hydroquinone as a sacrificial coreactant), whereas the cathode side is modified with copper. Idem in the case of (TMTSF)\(_2\)PF\(_6\) (c and d, respectively).
latter case, the absolute intensity of the photovoltage was found to be independent of the excitation side, confirming the symmetry breaking assigned to the Cu deposit (Figure 5d). Such an asymmetric photovoltage behavior was reported recently in the case of Janus graphene sheets and is observed here for the first time on a more complex Janus organic single crystal conductor. Finally, the comparison between a modified and an unmodified crystal confirms also that the chemical integrity of the Bechgaard salt was not altered by the copper electrodeposition process.

■ CONCLUSIONS

Using BPE has already proven to be successful for the fabrication of various asymmetric Janus particles. So far, the almost unique strategy was based on the site-selective electrodeposition of a redox-active precursor at one extremity of either carbon or metal bipolar electrodes. The choice of these substrates is straightforward because they are indeed very good conductors typically used as electrode materials for conventional electrochemistry. In an effort to extend the scope and potentials of BPE, Fabre and Bechgaard salts based on TMTTF and TMTSF moieties were selected and used as bipolar electrodes. This is the first demonstration of the use of organic crystal conductors in bipolar electrochemistry for which copper metal electrodeposition was chosen. The approach opens the route to new families of Janus particles that can be prepared by BPE in the bulk phase. The corresponding asymmetric objects were characterized by optical imaging and electron microscopy revealing that the edge of the crystal acting as an anode is very sensitive to electrodissolution. This drawback has been overcome by the addition of a sacrificial coreactant with an appropriate redox potential in order to carry out bipolar electrodeposition without altering the chemical structure of the crystal. Finally, asymmetric photovoltage generation could be observed as a result of the intrinsic break of symmetry provided by the site-selective metal deposition. We anticipate that this kind of new functional hybrid materials will readily be employed for applications in organic electronics and/or photocatalysis, especially for electronic display or photoactive electronic devices. Besides, more elaborated TTF-based conducting crystals with other functionalities will be considered as possible BPE organic electrodes.

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Notes
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